

(43) Date of publication:
 28.11.2001 Bulletin 2001/48

(51) Int. Cl. : C12Q 1/68, C07K 1/00,
 C07K 17/00, C07H 21/00

(21) Application number: 00915717.3

(86) International application number:
 PCT/US00/02490

(22) Date of filing: 01.02.2000

(87) International publication number:
 WO 00/47774 (17.08.2000 Gazette 2000/33)

(84) Designated Contracting States:
 AT BE CH CY DE DK ES FI FR GB GR IE IT LI LU
 MC NL PT SE
 Designated Extension States:
 AL LT LV MK RO SI

(30) Priority: 09.02.1999 US 246461

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(54) APTAMERS AS REAGENTS FOR HIGH THROUGHPUT SCREENING

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INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification: C12Q 1/68, C07H 21/00, C07K 1/00, C07K 17/00		A1	(11) International Publication Number: WO 00/47774 (43) International Publication Date: 17 August 2000 (17.08.2000)
(21) International Application Number: PCT/US00/02490			
(22) International Filing Date: 01 February 2000 (01.02.2000)		Published	
(30) Priority Data: 09/246,461 09 February 1999 (09.02.1999) US			
(60) Parent Application or Grant NEXSTAR PHARMACEUTICALS, INC. [/]; O. JANJIC, Nebojsa [/]; O. GOLD, Larry [/]; O. SWANSON, Barry, J. ; O.			
(54) Title: APTAMERS AS REAGENTS FOR HIGH THROUGHPUT SCREENING (54) Titre: APTAMERES UTILISES COMME REACTIFS DANS LA RECHERCHE A HAUT RENDEMENT			
(57) Abstract The invention relates to the use of aptamers in high throughput screening methods to determine whether a non-nucleic acid molecule, i.e. small molecule, binds to a target.			
(57) Abrégé La présente invention concerne l'utilisation d'aptamères dans des méthodes de recherche à haut rendement afin de déterminer si une molécule d'acide nucléique, c'est-à-dire une petite molécule, est fixée sur une cible.			

PCT

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<p>(51) International Patent Classification 7 : C12Q 1/68, C07K 1/00, 17/00, C07H 21/00</p>		<p>A1</p>	<p>(11) International Publication Number: WO 00/47774 (43) International Publication Date: 17 August 2000 (17.08.00)</p>
<p>(21) International Application Number: PCT/US00/02490 (22) International Filing Date: 1 February 2000 (01.02.00)</p>		<p>(81) Designated States: AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, DM, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SL, SZ, TZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).</p>	
<p>(30) Priority Data: 09/246,461 9 February 1999 (09.02.99) US</p>		<p>Published <i>With international search report. Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.</i></p>	
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<p>(54) Title: APTAMERS AS REAGENTS FOR HIGH THROUGHPUT SCREENING</p>			
<p>(57) Abstract</p>			
<p>The invention relates to the use of aptamers in high throughput screening methods to determine whether a non-nucleic acid molecule, i.e. small molecule, binds to a target.</p>			

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Description

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APTAMERS AS REAGENTS FOR HIGH THROUGHPUT SCREENING

FIELD OF THE INVENTION

10 5 This invention is directed to a high throughput screening (HTS) method that uses aptamers to facilitate the identification of non-aptamer ligands, typically small molecules. Aptamers can be identified that bind to a wide variety of targets and can be used to identify small molecules that can compete with the aptamers for binding to the target.

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BACKGROUND OF THE INVENTION

20 20 Drug discovery in most pharmaceutical companies is heavily focused on the identification of orally active compounds. High throughput screening (HTS) of appropriate libraries of compounds (generally small molecules) against validated targets constitutes one of the major activities in discovery research groups whose overall goal is to

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15 15 create and/or maintain a pipeline of new drug candidates. The development of HTS assays that allow efficient detection of active compounds from such libraries is thus a critically important component of drug discovery. With increasing numbers of both validated targets and compound libraries, HTS laboratories are under considerable pressure to find ways to increase throughput and lower cost. Assay development and validation is often 30 20 one of the bottlenecks in this process (Fox *et al.* (November 1998) *Drug Discovery & Development* (Supplement to R&D Magazine) pp. 32-37, incorporated by reference in its entirety).

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35 Aptamers (also termed nucleic acid ligands) are structurally unique nucleic acids capable of binding other molecules (i.e., targets) with high affinity and specificity.

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25 25 Aptamers are derived from large random libraries by the SELEX process in which iterative rounds of affinity selection and amplification are used to identify sequences with specific binding properties. The SELEX process is described in more detail below. To date, aptamers that bind to small organic molecules, carbohydrates, amino acids, peptides and 45 30 proteins have been identified, illustrating the multitude of binding specificities present in large collections of single stranded nucleic acid sequences (Gold *et al.* (1995) *Ann. Rev. Biochem.* **64**:763-797). Aptamers bind to protein targets including growth factors, enzymes, receptors and structural proteins in a highly specific manner and with 50 dissociation constants typically in the nanomolar (and sometimes picomolar) range. Given these unique binding properties, nuclease stabilized and appropriately formulated aptamers

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have substantial potential as therapeutics. For therapeutic uses, aptamers, like antibodies, are delivered parenterally because of their limited oral availability.

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The dogma for many years was that nucleic acids had primarily an informational role. Through a method known as Systematic Evolution of Ligands by EXponential

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enrichment, termed the SELEX process, it has become clear that nucleic acids have three dimensional structural diversity not unlike proteins. The SELEX process is a method for the *in vitro* evolution of nucleic acid molecules with highly specific binding to target molecules and is described in United States Patent Application Serial No. 07/536,428,

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filed June 11, 1990, entitled "Systematic Evolution of Ligands by EXponential Enrichment," now abandoned, United States Patent Application Serial No. 07/714,131,

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filed June 10, 1991, entitled "Nucleic Acid Ligands," now United States Patent No. 5,475,096, and United States Patent Application Serial No. 07/931,473, filed August 17, 1992, entitled "Methods for Identifying Nucleic Acid Ligands," now United States Patent

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No. 5,270,163 (see also WO 91/19813), each of which is specifically incorporated by reference herein. Each of these applications, collectively referred to herein as the SELEX Patent Applications, describes a fundamentally novel method for making a nucleic acid ligand to any desired target molecule. The SELEX process provides a class of products

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which are referred to as aptamers or nucleic acid ligands, each ligand having a unique sequence, and which has the property of binding specifically to a desired target compound or molecule. Each SELEX-identified nucleic acid ligand is a specific ligand of a given target compound or molecule. The SELEX process is based on the unique insight that nucleic acids have sufficient capacity for forming a variety of two- and three-dimensional structures and sufficient chemical versatility available within their monomers to act as ligands (form specific binding pairs) with virtually any chemical compound, whether monomeric or polymeric. Molecules of any size or composition can serve as targets.

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The SELEX method applied to the application of high affinity binding involves selection from a mixture of candidate oligonucleotides and step-wise iterations of binding, partitioning and amplification, using the same general selection scheme, to achieve virtually any desired criterion of binding affinity and selectivity. Starting from a mixture of nucleic acids, preferably comprising a segment of randomized sequence, the SELEX method includes steps of contacting the mixture with the target under conditions favorable for binding, partitioning unbound nucleic acids from those nucleic acids which have bound

5 specifically to target molecules, dissociating the nucleic acid-target complexes, amplifying
10 the nucleic acids dissociated from the nucleic acid-target complexes to yield a ligand-
15 enriched mixture of nucleic acids, then reiterating the steps of binding, partitioning,
dissociating and amplifying through as many cycles as desired to yield highly specific
20 5 high affinity nucleic acid ligands to the target molecule.

25 The SELEX method demonstrates that nucleic acids as chemical compounds can
15 form a wide array of shapes, sizes and configurations, and are capable of a far broader
repertoire of binding and other functions than those displayed by nucleic acids in
biological systems. SELEX or SELEX-like processes can be used to identify nucleic acids
10 which can facilitate any chosen reaction in a manner similar to that in which nucleic acid
20 ligands can be identified for any given target. In theory, within a candidate mixture of
approximately 10^{13} to 10^{18} nucleic acids, at least one nucleic acid exists with the
25 appropriate shape to facilitate each of a broad variety of physical and chemical
interactions.

30 15 The basic SELEX method has been modified to achieve a number of specific
objectives. For example, United States Patent Application Serial No. 07/960,093, filed
October 14, 1992, entitled "Method for Selecting Nucleic Acids on the Basis of Structure,"
35 now abandoned (see United States Patent No. 5,707,796) describes the use of the SELEX
process in conjunction with gel electrophoresis to select nucleic acid molecules with
20 specific structural characteristics, such as bent DNA. United States Patent Application
35 Serial No. 08/123,935, filed September 17, 1993, entitled "Photoselection of Nucleic Acid
Ligands," now abandoned (see United States Patent No. 5,763,177) describes a SELEX
40 based method for selecting nucleic acid ligands containing photoreactive groups capable of
binding and/or photocrosslinking to and/or photoactivating a target molecule. United
25 States Patent Application Serial No. 08/134,028, filed October 7, 1993, entitled "High-
Affinity Nucleic Acid Ligands That Discriminate Between Theophylline and Caffeine,"
now abandoned (see United States Patent No. 5,580,737), describes a method for
45 identifying highly specific nucleic acid ligands able to discriminate between closely
related molecules, which can be non-peptidic, termed Counter-SELEX. United States
30 Patent Application Serial No. 08/143,564, filed October 25, 1993, entitled "Systematic
Evolution of Ligands by EXponential Enrichment: Solution SELEX," now abandoned
50 (see United States Patent No. 5,567,588), describes a SELEX-based method which

5 achieves highly efficient partitioning between oligonucleotides having high and low
10 affinity for a target molecule.

15 The SELEX method encompasses the identification of high-affinity nucleic acid
ligands containing modified nucleotides conferring improved characteristics on the ligand,

20 5 such as improved in vivo stability or improved delivery characteristics. Examples of such
modifications include chemical substitutions at the ribose and/or phosphate and/or base
15 positions. SELEX process-identified nucleic acid ligands containing modified nucleotides
are described in United States Patent Application Serial No. 08/117,991, filed September
8, 1993, entitled "High Affinity Nucleic Acid Ligands Containing Modified Nucleotides,"
20 now abandoned (see United States Patent No. 5,660,985), that describes oligonucleotides
containing nucleotide derivatives chemically modified at the 5- and 2'-positions of
25 pyrimidines. United States Patent Application Serial No. 08/134,028, *supra*, describes
highly specific nucleic acid ligands containing one or more nucleotides modified with 2'-
amino (2'-NH₂), 2'-fluoro (2'-F), and/or 2'-O-methyl (2'-OMe). United States Patent
25 15 Application Serial No. 08/264,029, filed June 22, 1994, entitled "Novel Method of
Preparation of Known and Novel 2' Modified Nucleosides by Nucleophilic Displacement,"
describes oligonucleotides containing various 2'-modified pyrimidines.

30 The SELEX method encompasses combining selected oligonucleotides with other
selected oligonucleotides and non-oligonucleotide functional units as described in United

35 20 States Patent Application Serial No. 08/284,063, filed August 2, 1994, entitled
"Systematic Evolution of Ligands by EXponential Enrichment: Chimeric SELEX," now
United States Patent No. 5,637,459, and United States Patent Application Serial No.
40 25 08/234,997, filed April 28, 1994, entitled "Systematic Evolution of Ligands by
EXponential Enrichment: Blended SELEX," now United States Patent No. 5,683,867,
respectively. These applications allow the combination of the broad array of shapes and
45 other properties, and the efficient amplification and replication properties, of
oligonucleotides with the desirable properties of other molecules.

50 45 The SELEX method further encompasses combining selected nucleic acid ligands
with lipophilic compounds or non-immunogenic, high molecular weight compounds in a
30 diagnostic or therapeutic complex as described in United States Patent Application Serial
No. 08/434,465, filed May 4, 1995, entitled "Nucleic Acid Ligand Complexes." Each of

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the above described patent applications which describe modifications of the basic SELEX procedure are specifically incorporated by reference herein in their entirety.

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SUMMARY OF THE INVENTION

5 The present invention describes the use of aptamers to facilitate the identification of non-aptamer ligands. More specifically, the present invention includes the use of aptamers in competition binding assays to rapidly identify compounds that are capable of displacing the aptamers from their targets. The affinities of competitor compounds can be calculated from the known affinity of the aptamer for its target and the competition 10 profiles. The method is highly versatile and compatible with a variety of HTS platforms 20 since aptamers, as chemically synthesized molecules, can be labeled in a variety of ways without compromising their binding affinity.

25 The method of the invention is illustrated with two protein targets: platelet derived growth factor (PDGF) and wheat germ agglutinin (WGA). For each protein, a small 15 biased set of molecules is screened for their ability to displace the cognate aptamer: naphthalene sulfonic acid derivatives for PDGF and oligosaccharides for WGA. For both 30 PDGF and WGA, best ligands can be identified readily. Furthermore, binding affinities of the competitors correlate with their activities in *in vitro* assays (*infra*, and in United States Patent Number 5,780,222, issued July 14, 1998).

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BRIEF DESCRIPTION OF THE FIGURES

35 Figure 1 depicts the structures of small molecule oligoanion competitors used in the PDGF competition assay (dashed lines (c_{2v}) indicate a 2-fold axis of symmetry) along with the K_{dc} values calculated from the competition profiles according to equations (1)-(3) 40 and EC50 values derived from ^3T -thymidine uptake assays.

25 Figures 2A, 2B and 2C depict the competition profiles of the various small molecule oligoanion competitors for displacing the binding of a PDGF aptamer to PDGF.

45 Figure 3 depicts the structures of small molecule oligosaccharide competitors used in the WGA competition assay along with the K_{dc} values calculated from the competition 30 profiles according to equations (1)-(3).

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Figures 4A, 4B, 4C and 4D depict the competition profiles of the various small molecule oligosaccharide competitors for displacing the binding of the WGA aptamer to WGA.

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5 **DETAILED DESCRIPTION OF THE INVENTION**

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The present invention describes the use of aptamers as a new class of reagents for high throughput screening (HTS) that may be used in conjunction with or in place of cell-based assays, receptor binding and other protein-protein interaction assays. The identification of aptamers to protein targets generally requires 5-15 rounds of the SELEX process to achieve affinities in the nanomolar range. A round of the SELEX process using manual protocols takes about one day, and considerably less than that using automated protocols such as those described in United States Patent Application Serial No. 09/232,946, entitled "Method and Apparatus for the Automated Generation of Nucleic Acid Ligands," filed January 19, 1999, which is incorporated herein in its entirety.

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10 Aptamer identification is thus rapid.

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It is important to note that for use in competition assays described here, the affinity of aptamers for their targets need not be very high, since the affinity of first generation lead compounds will typically be considerably lower than that of an aptamer (e.g., in the micromolar range). For example, for the purposes of this invention it was useful to reduce the affinity of a previously identified PDGF aptamer (described in United States Patent No. 5,723,594, entitled "High Affinity PDGF Nucleic Acid Ligands," issued March 3, 1998, which is incorporated herein in its entirety) about 10-fold to facilitate the identification of competitors that bind PDGF with micromolar affinities. Lower affinity requirements may further speed up the identification of appropriate aptamers for competition assays. More generally, the affinities of aptamers and binding conditions (such as concentrations of the binding species) can be tuned to facilitate detection of ligands in a defined concentration range. Aptamers may be particularly useful for the identification of ligands to protein targets that do not have a known binding partner, such as orphan receptors.

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35 The use of competition binding screens with aptamers (or any other ligands) does not guarantee that all compounds from a library that are capable of binding to the target will be identified, since the assay requires competitive displacement. To decrease the

chance that active compounds may be missed, one can use more than one aptamer for the screens. It is important to note in this context, however, that in most cases, aptamers identified to protein targets generally compete with each other even when they belong to different sequence families or have different compositions (RNA, DNA or modified RNA). Nevertheless, given their size (typically 6-13 kDa), most aptamers are likely to cover a significant surface fraction of their protein targets which should reduce the problem of false negatives in competition assays. Furthermore, as illustrated with the WGA aptamers (described in United States Patent Number 5,780,228, entitled "High Affinity Nucleic Acid Ligands to Lectins," issued July 14, 1998, which is incorporated herein in its entirety), aptamers that bind to specific sites on proteins can be selected provided that a ligand that binds to that site is available.

High affinity binding of aptamers for protein targets is typically encoded in sequences of 20-40 nucleotides. The efficient encoding of high affinity binding allows aptamers to be synthesized entirely chemically, e.g. by the solid phase phosphoramidite method. Aside from the advantage of being able to control batch-to-batch variability and lower reagent cost, chemical synthesis facilitates the incorporation of various non-nucleic acid functionalities into aptamers in a manner that does not disrupt their exquisite binding properties. Therefore, although radiolabeled aptamers are used in the examples herein, aptamers can be labeled in a variety of other ways (e.g., with light-absorbing, fluorescent or chemiluminescent moieties, biotin, etc.) that may be more suitable for some HTS applications.

Definitions

Various terms are used herein to refer to aspects of the present invention. To aid in the clarification of the description of the components of this invention, the following definitions are provided.

As used herein, "aptamer" or "nucleic acid ligand" is a non-naturally occurring nucleic acid having a desirable action on a target. A desirable action includes, but is not limited to, binding of the target, catalytically changing the target, reacting with the target in a way which modifies/alters the target or the functional activity of the target, covalently attaching to the target as in a suicide inhibitor, facilitating the reaction between the target and another molecule. In the preferred embodiment, the action is specific binding affinity

5 for a target molecule, such target molecule being a three dimensional chemical structure
other than a polynucleotide that binds to the nucleic acid ligand through a mechanism
which predominantly depends on Watson/Crick base pairing or triple helix binding,
10 wherein the nucleic acid ligand is not a nucleic acid having the known physiological
15 function of being bound by the target molecule. Nucleic acid ligands include nucleic acids
that are identified from a candidate mixture of nucleic acids, said nucleic acid ligand being
a ligand of a given target, by the method comprising: a) contacting the candidate mixture
with the target, wherein nucleic acids having an increased affinity to the target relative to
the candidate mixture may be partitioned from the remainder of the candidate mixture; b)
10 partitioning the increased affinity nucleic acids from the remainder of the candidate
20 mixture; and c) amplifying the increased affinity nucleic acids to yield a ligand-enriched
mixture of nucleic acids.

25 As used herein, "non-aptamer ligands" or "non-nucleic acid molecule" is any
molecule that is not an aptamer. Typically this term includes but is not limited to small
25 molecules.

30 As used herein, "candidate mixture" is a mixture of nucleic acids of differing
sequence from which to select a desired aptamer. The source of a candidate mixture can
30 be from naturally-occurring nucleic acids or fragments thereof, chemically synthesized
nucleic acids, enzymatically synthesized nucleic acids or nucleic acids made by a
20 combination of the foregoing techniques. In a preferred embodiment, each nucleic acid
35 has fixed sequences surrounding a randomized region to facilitate the amplification
process.

40 As used herein, "nucleic acid" means either DNA, RNA, single-stranded or double-
stranded, and any chemical modifications thereof. Modifications include, but are not
40 limited to, those which provide other chemical groups that incorporate additional charge,
polarizability, hydrogen bonding, electrostatic interaction, and fluxionality to the nucleic
acid ligand bases or to the nucleic acid ligand as a whole. Such modifications include, but
45 are not limited to, 2'-position sugar modifications, 5-position pyrimidine modifications, 8-
position purine modifications, modifications at exocyclic amines, substitution of 4-
50 thiouridine, substitution of 5-bromo or 5-iodo-uracil; backbone modifications,
methylations, unusual base-pairing combinations such as the isobases isocytidine and

5 isoguanidine and the like. Modifications can also include 3' and 5' modifications such as
10 capping.

15 "SELEX" methodology involves the combination of selection of nucleic acid
ligands which interact with a target in a desirable manner, for example binding to a
20 protein, with amplification of those selected nucleic acids. Optional iterative cycling of
the selection/amplification steps allows selection of one or a small number of nucleic acids
which interact most strongly with the target from a pool which contains a very large
25 number of nucleic acids. Cycling of the selection/amplification procedure is continued
until a selected goal is achieved. The SELEX methodology is described in the SELEX
30 Patent Applications.

20 "Target" means any compound or molecule of interest for which a ligand is
desired. A target can be a protein, peptide, carbohydrate, polysaccharide, glycoprotein,
hormone, receptor, antigen, antibody, virus, substrate, metabolite, transition state analog,
25 cofactor, inhibitor, drug, dye, nutrient, growth factor, etc. without limitation.

30 15 As used herein, "solid support" is defined as any surface to which molecules may
be attached through either covalent or non-covalent bonds. This includes, but is not
35 limited to, membranes, plastics, magnetic beads, charged paper, nylon, Langmuir-Bodgett
40 films, functionalized glass, germanium, silicon, PTFE, polystyrene, gallium arsenide, gold
45 and silver. Any other material known in the art that is capable of having functional groups
50 such as amino, carboxyl, thiol or hydroxyl incorporated on its surface, is also
contemplated. This includes surfaces with any topology, including, but not limited to,
spherical surfaces and grooved surfaces.

40 "Partitioning" means any process whereby aptamers bound to target molecules can
be separated from nucleic acids not bound to target molecules. More broadly stated,
45 25 partitioning allows for the separation of all the nucleic acids in a candidate mixture into at
least two pools based on their relative affinity to the target molecule. Partitioning can be
accomplished by various methods known in the art. Nucleic acid-protein pairs can be
bound to nitrocellulose filters while unbound nucleic acids are not. Columns which
50 30 specifically retain nucleic acid-target complexes can be used for partitioning. For
example, oligonucleotides able to associate with a target molecule bound on a column
allow use of column chromatography for separating and isolating the highest affinity
nucleic acid ligands. Beads upon which target molecules are conjugated can also be used

to partition nucleic acid ligands in a mixture. Surface plasmon resonance technology can be used to partition nucleic acids in a mixture by immobilizing a target on a sensor chip and flowing the mixture over the chip, wherein those nucleic acids having affinity for the target can be bound to the target, and the remaining nucleic acids can be washed away.

5 Liquid-liquid partitioning can be used as well as filtration gel retardation, and density gradient centrifugation.

In its most basic form, the SELEX process may be defined by the following series of steps:

1) A candidate mixture of nucleic acids of differing sequence is prepared. The
10 candidate mixture generally includes regions of fixed sequences (i.e., each of the members
of the candidate mixture contains the same sequences in the same location) and regions of
randomized sequences. The fixed sequence regions are selected either: a) to assist in the
amplification steps described below; b) to mimic a sequence known to bind to the target;
or c) to enhance the concentration of a given structural arrangement of the nucleic acids in
15 the candidate mixture. The randomized sequences can be totally randomized (i.e., the
probability of finding a base at any position being one in four) or only partially
randomized (e.g., the probability of finding a base at any location can be selected at any
level between 0 and 100 percent).

2) The candidate mixture is contacted with the selected target under conditions favorable for binding between the target and members of the candidate mixture. Under these circumstances, the interaction between the target and the nucleic acids of the candidate mixture can be considered as forming nucleic acid-target pairs between the target and those nucleic acids having the strongest affinity for the target.

25 3) The nucleic acids with the highest affinity for the target are partitioned from those nucleic acids with lesser affinity to the target. Because only an extremely small number of sequences (and possibly only one molecule of nucleic acid) corresponding to the highest affinity nucleic acids exist in the candidate mixture, it is generally desirable to set the partitioning criteria so that a certain amount of the nucleic acids in the candidate mixture are retained during partitioning.

30 4) Those nucleic acids selected during partitioning as having relatively higher affinity to the target are then amplified to create a new candidate mixture that is enriched in nucleic acids having a relatively higher affinity for the target.

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5) By repeating the partitioning and amplifying steps above, the newly formed
candidate mixture contains fewer and fewer unique sequences, and the average degree of
affinity of the nucleic acids to the target will generally increase. Taken to its extreme, the
10 SELEX process will yield a candidate mixture containing one or a small number of unique
5 nucleic acids representing those nucleic acids from the original candidate mixture having
the highest affinity to the target molecule.

15 The SELEX Patent Applications describe and elaborate on this process in great
detail. Included are targets that can be used in the process; methods for the preparation of
the initial candidate mixture; methods for partitioning nucleic acids within a candidate
10 mixture; and methods for amplifying partitioned nucleic acids to generate enriched
20 candidate mixtures. The SELEX Patent Applications also describe ligand solutions
obtained to a number of target species, including protein targets wherein the protein is or is
not a nucleic acid binding protein.

25 In the preferred embodiment of this invention, aptamers are used in conjunction
15 with an existing HTS platform that allows inspection of multiple wells in an automated
format. Any and all HTS platforms are contemplated for use in the present invention. For
example, a target could be immobilized on a solid support and incubated with labeled
30 aptamer. Any labeling method is contemplated by the present invention, including but not
limited to radioactive, light-absorbing, fluorescent, chemiluminescent or other detectable
20 moieties. A library of non-aptamer candidate molecules could then be tested for their
ability to displace the aptamer from its target by measuring either the amount of labeled
35 aptamer displaced or the amount of labeled aptamer remaining on the solid support.
Alternatively, the aptamer could be immobilized on a solid support and incubated with a
labeled target. A library of non-aptamer candidate molecules could then be tested for their
40 ability to displace the labeled target as described above. Any known method for detecting
the displaced aptamer is contemplated by the present invention, including but limited to
direct detection or amplified detection such as that described in United States Application
45 Serial No. 09/157,206 filed September 18, 1998, entitled "Homogenous Detection of a
Target Through Nucleic Acid Ligand-Ligand Beacon Interaction," which is incorporated
30 herein in its entirety by reference.

50 In certain embodiments, the method can take place in solution with the aptamer
and small molecule competing simultaneously.

5

12

EXAMPLES

10 The Examples provided below are illustrative embodiments of the invention. They are not to be taken as limiting the scope of the invention.

5

Example 1. Material and MethodsMaterials

10 Human recombinant platelet derived growth factor, BB isoform (PDGF BB) was purchased from R&D Systems (Minneapolis, MN) as a carrier free lyophilized powder.

20 Wheat Germ (*Triticum vulgare*) Agglutinin (WGA) was from EY laboratories (San Mateo, CA).

25 Oligonucleotides were synthesized using an Applied Biosystems Model 394 oligonucleotide synthesizer according to optimized protocols. PDGF aptamer 20ta is a

synthetic 33-mer DNA oligo of sequence: 5'-CGGGCGCGTTCTCGTGGTTACTTTT
AGTCCCG (SEQ ID NO: 1), aptamer 20tb is a synthetic 27-mer DNA oligo of sequence:

30 5'-GGGCCGTTCCGGTTACTTTAGTCCC (SEQ ID NO: 2), and aptamer PD316 is a synthetic oligo containing some modified (2'-F and 2'-O-methyl, italic and bold letters,

35 respectively) bases, and an 18-atom PEG spacer replacing some bases, to increase serum stability, of sequence: 5'-T_{NH₂}CAGGUACG[PEG₁₈]CGTAGAGCAUCA[PEG₁₈]TGAT
CCUG-3'3'T (SEQ ID NO: 3). WGA aptamer 11.20 is a 98-mer RNA transcript with 2'-

40 aminopyrimidine bases of sequence: 5'-GGGAAAAGCGAAUCAUACACAAGAUUG
GUCGUACUGGACAGAGCCGUGGUAGAGGGAUUGGGACAAAGUGUCAGCUCC

45 25 GCCAGAGACCAACCGAGAA (SEQ ID NO: 4). PDGF aptamers were previously described in United States Patent No. 5,723,594, entitled "High Affinity PDGF Nucleic Acid Ligands," issued March 3, 1998. WGA aptamers were previously described in United States Patent No. 5,780,228, entitled "High Affinity Nucleic Acid Ligands to Lectins," issued July 14, 1998.

50 30 Oligoanions used in the PDGF aptamer competition assay were: Evans blue, trypan Blue, amaranth, sulfonazo III, New Coccine, *myoinositol* hexasulfate, SPADNS (2-(4-sulfophenylazo)-1,8-dihydroxy-3,6-naphthalene disulfonic acid), calcion and azocarmine B from Aldrich (Milwaukee, WI); NTSA (naphthalene 1,3,6-trisulfonic acid)

5

13

from Fluka, suramin from Calbiochem (La Jolla, CA) and sucrose octasulfate from Toronto Research Chemicals (Toronto, Canada).

10

Oligosaccharides used in the WGA aptamer competition assay were: N-acetyl-D-glucosamine (GlcNAc) from Sigma (St. Louis, MO); N,N'-diacetylchitobiose (GlcNAc)₂,

15

5 N,N',N"-triacetylchitotriose (GlcNAc)₃, N,N',N",N""-tetraacetyltetraose (GlcNAc)₄, galactose 1- β -3(fucose- α -4)glucosamine (Lewis A trisaccharide), galactose 1- β -4(fucose 1- α -3)glucosamine (Lewis X trisaccharide), fucose1- α -2 galactose1- β -4(fucose 1- α -3)glucosamine (Lewis Y tetrasaccharide), Sialyl Lewis A, Sialyl Lewis X, fucose1- α -4glucosamine, fucose1- α -3glucosamine, N-acetyl-lactosamine (LacNAc), 3' α -Sialyl-N-acetyllactosamine (α -Sialyl-LacNAc), 3' β -Sialyl-N-acetyllactosamine (β -Sialyl-LacNAc), from Toronto Research Chemicals (Toronto, Canada).

20

Methods

Small molecule/aptamer competition assays

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15 Small molecule oligoanions were screened for their ability to compete with PDGF aptamer 20tb for binding to the PDGF-BB protein. The small molecule (or cold 20tb) competitors at varying concentrations were mixed with the 5'-³²P end-labeled 20tb ligand at room temperature. PDGF was then added to the mixture and allowed to equilibrate for 20 45 minutes at room temperature followed by 15 minutes at 37°C. The competition mixtures (90 μ L volume) all had ³²P end-labeled 20tb at 1.16 nM, PDGF-BB at 1 nM and were in 25 mM Hepes, 137 mM NaCl, 2.7 mM KCl, 1 mM MgCl₂, 1 mM CaCl₂, 0.067% 35 human serum albumin (HSA), at pH 7.4 (HBSMC). Aptamer bound to PDGF was separated from free aptamer by filtration through pre-washed 0.45 μ M nitrocellulose 25 membrane filters (Millipore; Bedford, MA). Filters were washed with 5 mL HBSMC at 40 room temperature. The filters were placed in tubes with scintillant and the radioactivity counted to quantitate the fraction of ³²P end-labeled 20tb bound at each competitor concentration.

45

Small molecule oligosaccharides were screened for their ability to compete with 30 WGA aptamer 11.20 for binding to wheat germ agglutinin protein. The small molecule (or cold 11.20) competitors at varying concentrations were mixed with the 5'-³²P end-labeled 11.20 ligand at room temperature. Then WGA was added to the mixture and 50 allowed to equilibrate for 60 minutes at room temperature. The competition mixtures (90

Analysis of competition data

10 Competition experiments were analyzed using equation (1) to determine the
20 concentration of free protein, $[P_F]$, as a function of the total concentration of competitor
added, $[C_T]$:

$$[P_E] = [P_T] / (1 + K_{dp} [R_T] / (1 + K_{dp} [P_E]) + K_{dc} [C_T] / (1 + K_{dc} [P_E])) \quad (1)$$

25 where K_{dR} is the binding constant of species R to the protein (assuming 1:1 stoichiometry) and K_{dC} is the binding constant of species C, the competitor, to the protein (assuming 1:1 stoichiometry). Equation (1) was solved to self-consistency of $[P_F]$ to a precision of 1×10^{-15} . Using these values of $[P_F]$, the concentration of protein-ligand complex $[PR]$ as a
 30 function of $[C_T]$ was determined using equation (2):
 20

$$[PR] = K_{PR} [R_T] [P_F] (1 + K_{PF} [P_F]) \quad (2)$$

40 Since the experimental data are expressed as %[PR], the calculated concentration of [PR]
 25 was normalized by $[PR_0]$, the [PR] in the absence of the competitor. $[PR_0]$ was obtained by
 solving equation (1) where $[C_T] = 0$. The maximum (M) and minimum (B) %[PR] were
 allowed to float during the analysis as shown in equation (3):

$$\%[\text{PR}] = [\text{PR}] / [\text{PR}_0] (M-B) + B \quad (3)$$

50 A non-linear least-squares fitting procedure was used as described by Bates and Watts (Bates and Watts (1988) in *Non-Linear Regression Analysis and its Applications*, D M

5 Bates and D.G. Watts editors, John Wiley & Sons, NY, NY). The program used was
originally written in C program language by Dominic Zichi and Brenda Javornic, NeXstar
10 Pharmaceuticals, Inc. The data were fit to equations (1) to (3) to obtain best fit parameters
for K_{dC} , M and B as a function of $[C_T]$ while leaving K_{dR} and $[P_T]$ constant.

5

Inhibition of PDGF stimulated [³H]-thymidine incorporation

15 A 10 rat smooth muscle cells were plated in 96 well tissue culture plates at $7-8 \times 10^3$
per well in 100 μ L DME + 4.5 g/L glucose + HEPES + 0.1% FBS and starved for about
20 hours. Aptamers or small molecule oligoanions were titrated across the 96 well plate in
10 triplicate wells from 1 mM to 0.3 μ M by 1:3 dilution steps for the oligoanions and from 1
20 μ M to 0.3 nM by 1:3 steps for the aptamers. Immediately, PDGF-BB was added to 10
ng/mL to all wells except the unstimulated controls. Positive controls had only PDGF-BB,
no other compounds. After 6 hours at 37°C in 5% CO₂, ³H-thymidine was added (0.25
25 μ Ci per well) and incubated for another 24 hours. Cells were lysed in 1% triton-X 100 for
15 20-30 minutes on a slow shaker at room temperature, then harvested onto 96 well glass
fiber filter plates (Packard) and dried. Scintillant was added and incorporated ³H-
thymidine radioactivity counted. EC₅₀ values for inhibition of PDGF-BB-induced ³H-
30 thymidine uptake were obtained by fitting the data to the nonlinear regression with
variable slope model of the GraphPad Prism computer program (GraphPad Software).

20

35 Example 2. PDGF aptamer competition assay

For the PDGF competition assay, a DNA PDGF aptamer sequence identified
previously was used. The aptamer and methods used herein are described in detail in
40 Example One. Aptamers obtained in this experiment bind preferentially to the B-chain of
PDGF in a manner that inhibits receptor binding and PDGF-BB-induced DNA synthesis *in*
vitro. Using photo-crosslinking experiments, it has been shown that a specific nucleotide
45 in the aptamer interacts with phenylalanine-84 of the PDGF B-chain which is located near
the region of PDGF known to be involved in receptor binding (Green *et al.* (1996))
50 Biochemistry 35:14413-24). To facilitate detection of competitors in the micromolar
range, the affinity of the PDGF aptamer 20ta ($K_d=50$ pM), a 33-mer, was deliberately
reduced about 10-fold by additional truncations. The aptamer used for competition
binding experiments, 20tb, is a 27-mer that binds to PDGF-BB with a K_d of 0.5 nM.

5 A panel of oligoanions, mostly in the naphthalene sulfonic acid class, were
10 screened for their ability to displace a ^{32}P -radiolabeled aptamer (20tb) from PDGF-BB.
15 These compounds were chosen because suramin and several other members of this family
inhibit the binding of PDGF to cells that express the PDGF receptor (Garrett *et al.* (1984)
20 5 PNAS 81:7466-7470; Powis *et al.* (1992) Cancer Chemother. Pharmacol. 31:223-228).
25 Since the PDGF aptamer also inhibits receptor binding (and with considerably greater
potency), it was reasonable to expect that suramin and the aptamer bind to PDGF in a
30 mutually exclusive manner. This was a deliberately biased library and was not intended to
approximate any conventional or combinatorial libraries but simply to demonstrate that
35 10 such competition assays are feasible. A nitrocellulose filter binding method was used to
separate bound from unbound aptamer. Structures of competitors used in this competition
assay are shown in Figure 1 along with the K_{dC} values calculated from the competition
40 15 profiles (Figure 2) according to equations (1)-(3) (Experimental Methods). For all
competition experiments, the concentration of the ^{32}P end-labeled 20tb aptamer and
45 20 PDGF-BB was 1.16 nM and 1.0 nM, respectively. Binding reactions were done at 37°C
and the time of incubations was at least 60 minutes to insure that the equilibrium was
established ($t_{1/2}$ of dissociation for the aptamer from PDGF-BB is about 2 min.). Fast
dissociation kinetics are clearly advantageous in these assays since they reduce the length
50 25 of time required to reach equilibrium.
20 Among the eleven compounds in the naphthalene sulfonic acid derivatives class,
35 there is clearly a range of affinities for PDGF. Suramin, a hexaanion, is actually not the
best ligand for PDGF-BB. Inspection of this set clearly suggests that the placement of the
40 30 sulfonic acid groups (or anions in general) in space is a strong determinant of binding
affinity. For example, naphthalene 1,3,6-trisulfonic acid binds with a K_{dC} of 870 μM
45 25 whereas SPADNS (another trianion with certain structural similarity to naphthalene 1,3,6-
trisulfonic acid) binds with a K_{dC} of 19 μM . The total number of negative charges seems
50 35 to be less important than their appropriate placement (compare suramin, a hexaanion with
myoinositol hexasulfate, also a hexaanion or sucrose octasulfate, an octaanion).

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Example 3. Effect of ligands on PDGF-induced 3 T-thymidine synthesis in A10 rat smooth muscle cells

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The same panel of small molecule oligoanions was tested for its effect on PDGF-BB-induced 3 T-thymidine incorporation in A10 rat smooth muscle cells as described in Example 1. The EC50 values for all compounds tested were calculated as described in Example 1 and are listed in Figure 1.

15

Example 4. Wheat Germ Agglutinin aptamer competition assay

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For the WGA competition assay, a 2'-aminopyrimidine RNA aptamer 11.20 identified previously was used. The aptamer and methods used herein are described in detail in Example 1. Aptamer 11.20 was selected by incubating a randomized nucleic acid library with WGA, removing the unbound molecules and then displacing the aptamers bound to a specific site with a competitor, (GlcNAc)₃. Thus, unlike the PDGF aptamer, which was selected for high affinity binding anywhere on the protein, the WGA aptamers were selected for binding to a specific site, the (GlcNAc)₃ binding site. Aptamer 11.20 and related aptamers isolated by this procedure potently inhibited WGA-mediated agglutination of sheep erythrocytes. Not surprisingly, aptamer 11.20 and related aptamers could be displaced with (GlcNAc)₃ (as described in United States Patent No. 5,780,228, entitled, "High Affinity Nucleic Acid Ligands to Lectins," issued July 14, 1998).

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A group of carbohydrates related to GlcNAc, were tested for their ability to displace radiolabeled aptamer 11.20 from WGA. As with PDGF, nitrocellulose filter binding assay was used to separate bound from unbound aptamer. Structures of competitors used in this assay are shown in Figure 3 along with the K_{dC} values calculated from the competition profiles (Figure 4) according to equations (1)-(3). Binding conditions are described in detail in Example 1.

45

Among the carbohydrates tested there is a wide range of affinities for WGA. The best ligands were (GlcNAc)₄, (GlcNAc)₃, and (GlcNAc)₂, in that order (Figures 3 and 4). This result is in agreement with previous observations (Goldstein and Poretz (1986) in: The Lectins. Properties, Functions, and Applications in Biology and Medicine, Academic Press, NY, pp 233-247.). GlcNAc was not a competitor in this concentration range. The ability of (GlcNAc)₃ and GlcNAc to inhibit WGA-mediated agglutination was tested previously (see United States Patent No. 5,780,228, entitled "High Affinity Nucleic Acid

10 in a functional assay.

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Claims

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What is claimed is:

10 1. A method for determining whether a non-nucleic acid molecule binds to a target comprising displacing a nucleic acid ligand having an affinity for said target with
5 said non-nucleic acid molecule.

15 2. A method for determining whether a non-nucleic acid molecule binds to a target comprising:

- 10 a) immobilizing a target to a solid support;
- 20 b) contacting a labeled nucleic acid ligand with said target wherein binding of the nucleic acid ligand to the target occurs to form a complex;
- 25 c) adding a non-nucleic acid molecule to the complex wherein a non-nucleic acid molecule that competes with the nucleic acid ligand for binding to the target will disrupt the complex and liberate the nucleic acid ligand; and
- 30 d) detecting the liberated nucleic acid ligand thereby determining that said non-nucleic acid molecule binds to said target.

35 3. The method of claim 2 wherein said nucleic acid ligand is identified by the method comprising:

- 20 i) preparing a candidate mixture of nucleic acids;
- 35 ii) contacting the candidate mixture of nucleic acids with said target, wherein nucleic acids having an increased affinity to said target relative to the candidate mixture may be partitioned from the remainder of the candidate mixture;
- 40 iii) partitioning the increased affinity nucleic acids from the remainder of the candidate mixture; and
- 45 iv) amplifying the increased affinity nucleic acids to yield a mixture of nucleic acids enriched for nucleic acids with relatively higher affinity and specificity for binding to said target, whereby a nucleic acid ligand may be identified.

50 30 4. The method of claim 2 wherein said label is selected from the group consisting of radio activity, fluorescence, chemiluminescence.

5

20
5. The method of claim 2 wherein said non-nucleic acid molecule is a small molecule.

10

6. A method for determining whether a non-nucleic acid molecule binds to a target comprising:

15

- a) immobilizing a nucleic acid ligand to a solid support;
- b) contacting a labeled target with said nucleic acid ligand wherein binding of the nucleic acid ligand to the target occurs to form a complex;
- c) adding a non-nucleic acid molecule to the complex wherein a non-nucleic acid molecule that competes with the nucleic acid ligand for binding to the target will disrupt the complex and liberate the target; and
- d) detecting the liberated target thereby determining that said non-nucleic acid molecule binds to said target.

20

25
15. The method of claim 6 wherein said nucleic acid ligand is identified by the method comprising:

30

- i) preparing a candidate mixture of nucleic acids;
- ii) contacting the candidate mixture of nucleic acids with said target, wherein nucleic acids having an increased affinity to said target relative to the candidate mixture may be partitioned from the remainder of the candidate mixture;
- iii) partitioning the increased affinity nucleic acids from the remainder of the candidate mixture; and
- iv) amplifying the increased affinity nucleic acids to yield a mixture of nucleic acids enriched for nucleic acids with relatively higher affinity and specificity for binding to said target, whereby a nucleic acid ligand may be identified.

40

45
8. The method of claim 6 wherein said label is selected from the group consisting of radio activity, fluorescence, chemiluminescence.

50

30
9. The method of claim 6 wherein said non-nucleic acid molecule is a small molecule.

<u>Competitor</u>	<u>Structure</u>	<u>K_{dC}, μM</u>	<u>$EC50$, μM</u>
Evans Blue		0.15 ± 0.04	5.6
Trypan Blue		0.26 ± 0.06	6.1
Suramin		1.1 ± 0.4	57
Calcion		7.8 ± 1.2	280
SPADNS		19 ± 6	100
Azocarmine B		22 ± 3.0	>1000

Figure 1

2/10

<u>Competitor</u>	<u>Structure</u>	<u>K_{dC}, μM</u>	<u>$EC50$, μM</u>
Amaranth		48 ± 16	510
Sulfonazo III		74 ± 27	>1000
New Coccine		120 ± 20	930
<i>myoinositol</i> hexasulfate		720 ± 200	>1000
Naphthalene 1,3,6-trisulfonic acid		870 ± 400	>1000
Sucrose octasulfate		1740 ± 450	>1000

Figure 1, continued

<u>Competitor</u>	<u>Structure</u>	<u>K_{dc}, μM</u>	<u>EC50, μM</u>
PDGF Aptamer (20ta)		0.000050 ± 0.000055	
PDGF Aptamer (20tb)		0.00039 ± 0.00016	
PDGF Aptamer(PD 316)		0.000070 ± 0.000030	0.005
VEGF Aptamer (31838SP)		>1	>1

Figure 1, continued

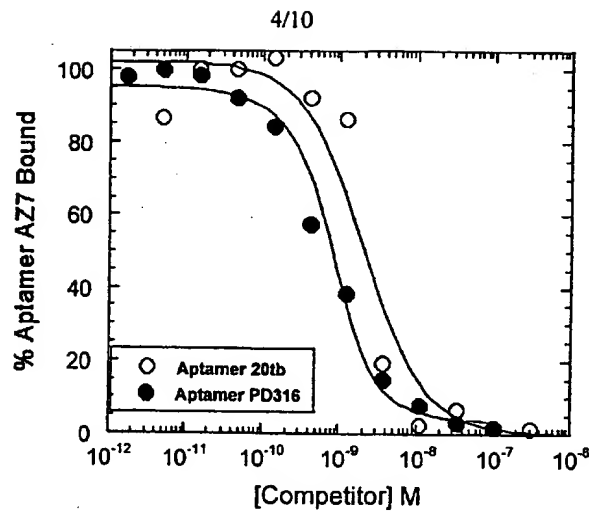


FIGURE 2A

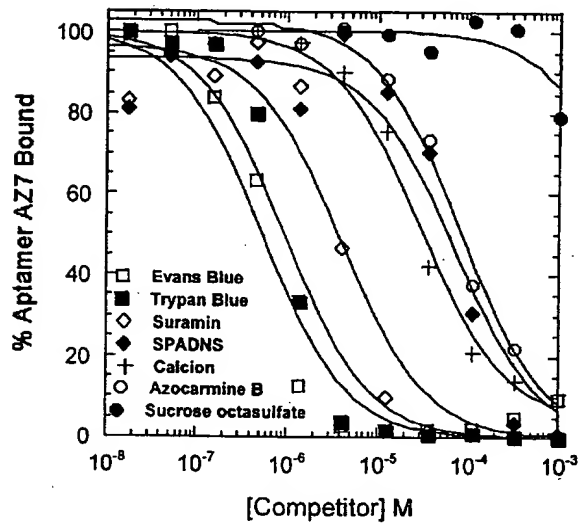


FIGURE 2B

5/10

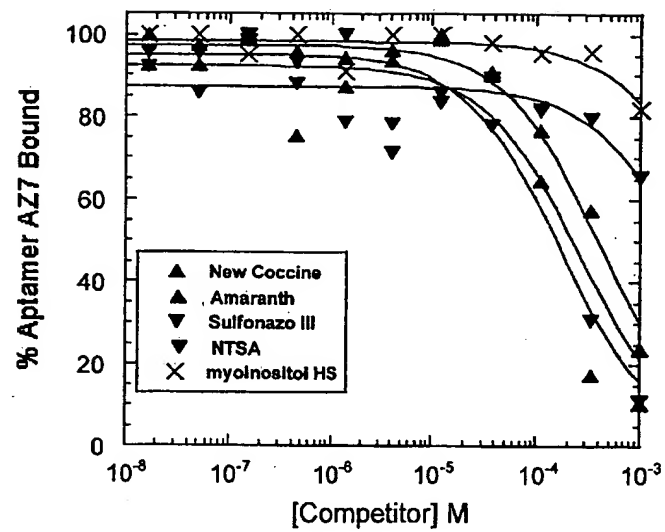


FIGURE 2C

<u>Competitor</u>	<u>Structure</u>	<u>K_{dC}, μM</u>
GlcNAc		>10000
$(\text{GlcNAc})_2$		83 ± 21
$(\text{GlcNAc})_3$		4 ± 0.3
$(\text{GlcNAc})_4$		1.3 ± 0.3
Fuc1- α -4GlcNAc		8100 ± 5200
Fuc1- α -3GlcNAc		6600 ± 5300

Figure 3

<u>Competitor</u>	<u>Structure</u>	<u>K_{dC}, μM</u>
Lewis A trisaccharide {Gal1-B-3[Fuc1- α -4] GlcNAc}		1500 \pm 400
Lewis X trisaccharide {Gal1-B-4[Fuc1- α -3] GlcNAc}		2600 \pm 1400
Lewis Y tetrasaccharide {Fuc1- α -2Gal1-B-4 [Fuc1- α -3]GlcNAc}		5400 \pm 7000
Sialyl Lewis A		6100 \pm 1000
Sialyl Lewis X		2800 \pm 300

Figure 3, continued

<u>Competitor</u>	<u>Structure</u>	<u>K_{dC}, mM</u>
LacNAc (Gal1-B-4GlcNAc)		480 ± 250
Sialyl- α -3LacNAc		540 ± 170
Sialyl- β -3LacNAc		440 ± 200

Figure 3, continued

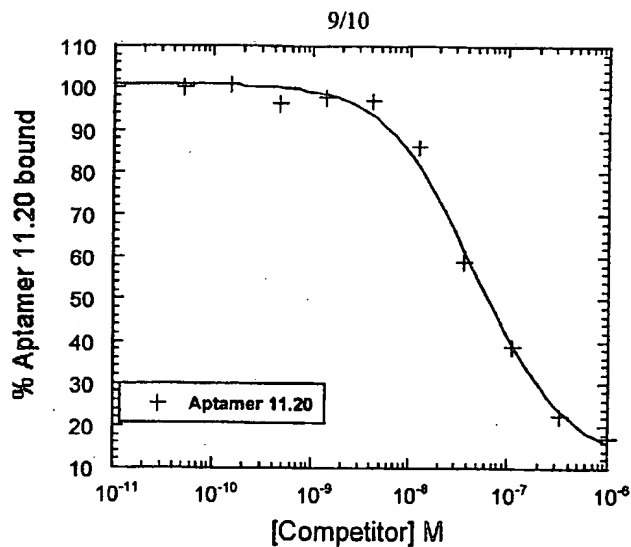


FIGURE 4A

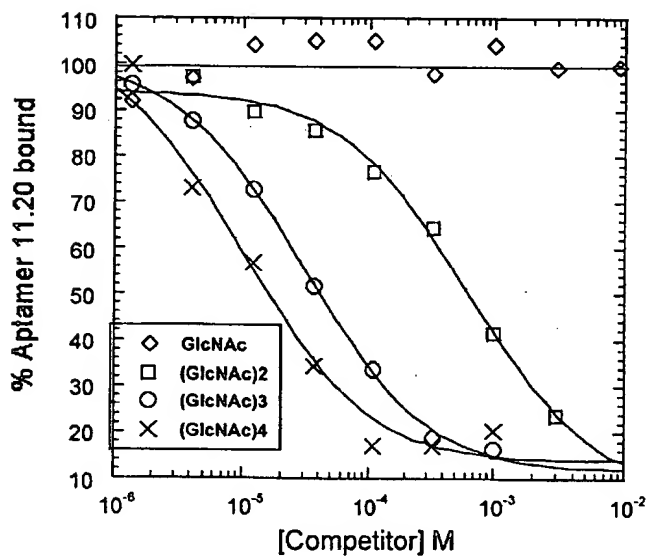


FIGURE 4B

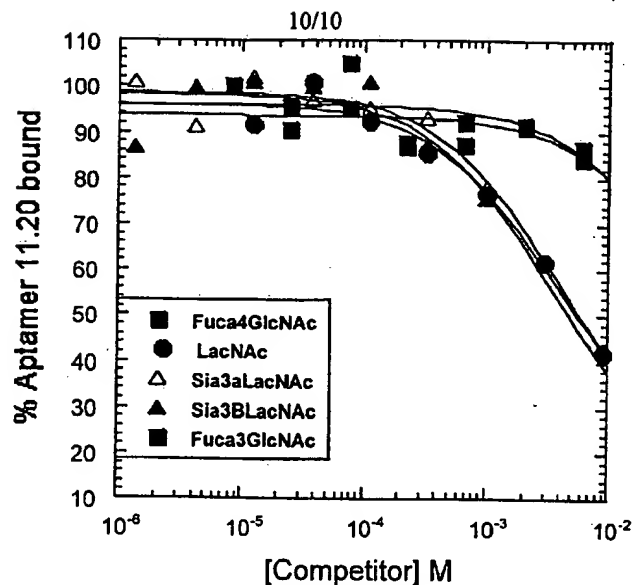


FIGURE 4C

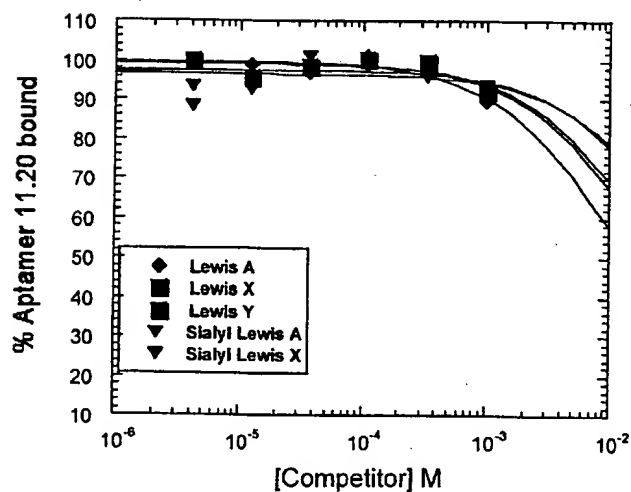


FIGURE 4D

SEQUENCE LISTING

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The a's and g's at positions 10, 15, 17, 22 and 30
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INTERNATIONAL SEARCH REPORT

International application No.
PCT/US00/02490

A. CLASSIFICATION OF SUBJECT MATTER IPC(7) C12Q 1/68; C07K 1/00, 17/00; C07H 21/00 US CL 435/6; 530/402, 413, 810, 812; 536/23.1 According to International Patent Classification (IPC) or to both national classification and IPC																
B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) U.S. : 435/6; 530/402, 413, 810, 812; 536/23.1																
Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched																
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) WEST (U.S. PATENTS) search terms: competitive, binding, displacing, nucleic acid, ligand, target, support																
C. DOCUMENTS CONSIDERED TO BE RELEVANT <table border="1" style="width: 100%; border-collapse: collapse;"> <thead> <tr> <th style="text-align: left; padding: 2px;">Category*</th> <th style="text-align: left; padding: 2px;">Citation of document, with indication, where appropriate, of the relevant passages</th> <th style="text-align: left; padding: 2px;">Relevant to claim No.</th> </tr> </thead> <tbody> <tr> <td style="text-align: center; padding: 2px;">X</td> <td style="padding: 2px;">US 5,780,228 A (PARMA et al.) 14 July 1998, entire document, especially paragraph bridging columns 18 and 19.</td> <td style="text-align: center; padding: 2px;">1</td> </tr> <tr> <td style="text-align: center; padding: 2px;">Y</td> <td></td> <td style="text-align: center; padding: 2px;">2-9</td> </tr> <tr> <td style="text-align: center; padding: 2px;">X</td> <td style="padding: 2px;">US 5,580,737 A (POLISKY et al.) 03 December 1996, entire document, especially column 17, lines 49-62, and column 18, lines 48-60.</td> <td style="text-align: center; padding: 2px;">1</td> </tr> <tr> <td style="text-align: center; padding: 2px;">Y</td> <td style="padding: 2px;">US 5,789,163 A (DROLET et al.) 04 August 1998, entire document, especially column 4, lines 19-37, column 3, lines 44-46, and column 15, lines 57-58.</td> <td style="text-align: center; padding: 2px;">2-9</td> </tr> </tbody> </table>		Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.	X	US 5,780,228 A (PARMA et al.) 14 July 1998, entire document, especially paragraph bridging columns 18 and 19.	1	Y		2-9	X	US 5,580,737 A (POLISKY et al.) 03 December 1996, entire document, especially column 17, lines 49-62, and column 18, lines 48-60.	1	Y	US 5,789,163 A (DROLET et al.) 04 August 1998, entire document, especially column 4, lines 19-37, column 3, lines 44-46, and column 15, lines 57-58.	2-9
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Y	US 5,789,163 A (DROLET et al.) 04 August 1998, entire document, especially column 4, lines 19-37, column 3, lines 44-46, and column 15, lines 57-58.	2-9														
<input type="checkbox"/> Further documents are listed in the continuation of Box C. <input type="checkbox"/> See patent family annex.																
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